

Durham Research Online

Deposited in DRO:

15 January 2018

Version of attached file:

Accepted Version

Peer-review status of attached file:

Peer-reviewed

Citation for published item:

Williams, Samual T. and Maree, Naudene and Taylor, Peter and Belmain, Steven R. and Keith, Mark and Swanepoel, Lourens H. (2018) 'Predation by small mammalian carnivores in rural agro-ecosystems : an undervalued ecosystem service?', *Ecosystem services*, 30 (C). 362–371.

Further information on publisher's website:

<https://doi.org/10.1016/j.ecoser.2017.12.006>

Publisher's copyright statement:

© 2018 This manuscript version is made available under the CC-BY-NC-ND 4.0 license
<http://creativecommons.org/licenses/by-nc-nd/4.0/>

Additional information:

Use policy

The full-text may be used and/or reproduced, and given to third parties in any format or medium, without prior permission or charge, for personal research or study, educational, or not-for-profit purposes provided that:

- a full bibliographic reference is made to the original source
- a [link](#) is made to the metadata record in DRO
- the full-text is not changed in any way

The full-text must not be sold in any format or medium without the formal permission of the copyright holders.

Please consult the [full DRO policy](#) for further details.

Williams, S.T., Maree, N., Taylor P, Belmain, S.R., Keith, M., Swanepoel, L.H. (2017)
Predation by small mammalian carnivores in rural agro-ecosystems: An undervalued
ecosystem service? Ecosystem Services. DOI: [10.1016/j.ecoser.2017.12.006](https://doi.org/10.1016/j.ecoser.2017.12.006).

Predation by small mammalian carnivores in rural agro- ecosystems: An undervalued ecosystem service?

Samual T. Williams^{a,b}, Naudene Maree^a, Peter Taylor^{c,d}, Steven R. Belmain^e, Mark Keith^f,
Lourens H. Swanepoel^a

^a Department of Zoology, School of Mathematical & Natural Sciences, University of Venda,
Thohoyandou, South Africa

^b Department of Anthropology, Durham University, Durham, United Kingdom

^c South African Research Chair on Biodiversity Value & Change, University of Venda,
Thohoyandou, South Africa

^d School of Life Sciences, University of KwaZulu-Natal, Private Bag X54001, Durban 4000,
South Africa

^e Natural Resources Institute, University of Greenwich, Chatham Maritime, Kent, United
Kingdom

^f Eugène Marais Chair of Wildlife Management, Mammal Research Institute, University of
Pretoria, 0002, South Africa

24

25 Abstract

26 Africa is endowed with a diverse guild of small carnivores, which could benefit stakeholders
27 by providing ecosystem services while fostering conservation tolerance for carnivores. To
28 investigate the potential of small carnivores for the biological control of rodents within agro-
29 ecosystems, we assessed both the ecological and social landscapes within two rural villages in
30 the Vhembe Biosphere Reserve, South Africa. We employed a camera trapping survey
31 underpinned by an occupancy modelling framework to distinguish between ecological and
32 observation processes affecting small carnivore occupancy. We also used questionnaires to
33 investigate perceptions of small carnivores and their role in pest control. We found the greatest
34 diversity of small carnivores in land used for cropping in comparison to grazing or settlements.
35 Probability of use by small carnivores was influenced negatively by the relative abundance of
36 domestic dogs and positively by the relative abundance of livestock. Greater carnivore diversity
37 and probability of use could be mediated through habitat heterogeneity, food abundance, or
38 reduced competition from domestic carnivores. Village residents failed to appreciate the role
39 of small carnivores in rodent control. Our results suggest that there is significant, although
40 undervalued, potential for small carnivores to provide ecosystem services in agro-ecosystems.

41

42 1. Introduction

43 Rodents cause significant damage to crops in small-holder farms in Africa (Granjon and
44 Duplantier, 2009; Monadjem *et al.*, 2015; Singleton, 2010; Swanepoel *et al.*, 2017). Existing
45 rodent control is highly reactive and almost exclusively based on the use of rodenticides. This
46 heavy reliance on poisons has led to increasing problems with the development of behavioural
47 and physiological resistance, environmental contamination, and non-target poisoning (Buckle
48 and Smith, 2015). Ecologically-based rodent management (EBRM) is a term popularised more
49 than 20 years ago (Singleton *et al.*, 1999) with an aim to re-emphasize the importance of
50 understanding rodent biology and behaviour of different species as well as agro-ecological and
51 socio-economic contexts. While traditional rodent pest solutions emphasized over-reliance on

poisons, EBRM advocates less harmful and sustainable solutions such as biological control through increasing ecosystem services of natural predation for pest control. Several studies have shown that the adoption of EBRM strategies for rodent pest management can be highly effective in reducing rodent damage whilst reducing farmer reliance on rodenticides (Brown *et al.*, 2006; Jacob *et al.*, 2010). EBRM has recently gained traction in small-holder agro-ecosystems in Africa (Massawe *et al.*, 2011; Monadjem *et al.*, 2015; Taylor *et al.*, 2012).

In smallholder agro-ecosystems, and many other modified landscapes, the removal of apex carnivore species from most human inhabited areas of Africa may have facilitated increased mesocarnivore abundance (Caro and Stoner, 2003; Prugh *et al.*, 2009; Ritchie and Johnson, 2009). Such increases might cause several ecological services or disservices to human communities. For example, small carnivores such as the red fox (*Vulpes vulpes*) provide valuable ecosystem services such as seed dispersal and potentially controlling populations of small mammals, regulating their impacts on keystone plant species and threatened habitats in Europe (Cancio *et al.*, 2017). In contrast, in Africa the importance of small carnivores around small-holder farming systems is well-recognised in terms of human-wildlife conflict and ecosystem disservices (Blaum *et al.*, 2009; Gusset *et al.*, 2009; Woodroffe *et al.*, 2005), but is less understood in terms of potential ecosystem services (Roemer *et al.*, 2009). This is unfortunate as Africa has a rich small carnivore assemblage, which could provide key ecosystem services to surrounding communities (Schuette *et al.*, 2013). Furthermore, the relatively large number of small-sized farms and small settlement areas in sub-Saharan Africa (Lowder *et al.*, 2016) are interspersed within a mosaic of semi-natural habitat that can increase human-wildlife conflict (Crooks, 2002; Lamarque *et al.*, 2009). As farm sizes in Africa are likely to continue to decline and further fragment the landscape (Masters *et al.*, 2013), there is a real risk of further natural habitat loss, trophic collapse and loss of potential ecosystem services provided by small carnivores (Dobson *et al.*, 2006).

Although the use of biological control is well established for many insect pests in agricultural production (Vincent *et al.*, 2007), it is not yet commonplace for rodent pests. The potential of

avian predators to provide ecosystem services for the control of pest rodents has been recently reviewed (Labuschagne *et al.*, 2016), highlighting that some species, such as barn owls (*Tyto alba*), are able to control rodent pests in some in agricultural contexts. Recent research suggests that domestic cats and dogs may increase the landscape of fear around rural homesteads, resulting in lower rates of rodent activity and food intake (Mahlaba *et al.*, 2017). This indirect mechanism, affecting rodent behaviour, could work synergistically with direct control mechanisms such as predation of rodents by domestic carnivores, which could reduce rodent density (Krijger *et al.*, 2017). Little attention, however, has been given to the potential services or disservices of wild terrestrial carnivores in terms of rodent pest control.

Thus, the first objective of our study was to understand which small- and medium-sized mammalian carnivores (< 15 kg, hereafter referred to as small carnivores) were present in and around rural farming communities in the study area. Secondly, we set out to determine the influence of the abundance of domestic animals (livestock and pets) on the probability of use of an area by small carnivores; and also assess how the species richness of the small carnivore community was influenced by land use. Thirdly, we wanted to capture the knowledge and opinions of smallholder farming communities with respect to small carnivores. This will provide an initial yet essential step towards understanding the potential ecosystem services provided by small carnivores in rural agro-ecosystems, to help inform the development of EBRM strategies with a strengthened biological control component.

2. Methods

2.1. Study area

We conducted the study at two rural sites (Ka-Ndengeza: S23.31003° E30.40981° and Vyeboom: S23.15174° E30.39278°) in the Vhembe Biosphere Reserve, South Africa (Appendix S1). Both sites receive an annual rainfall of 700-800 mm per year, with a hot wet season from October to March and a cool dry season from May to August (Hijmans *et al.*, 2005). Natural vegetation is classified as Granite Lowveld and Gravelotte rocky bushveld

(Mucina and Rutherford, 2006). Vegetation is characterised by tall shrubs with few trees to moderately dense low woodland on the deep sandy uplands dominated by *Combretum zeyheri* and *C. apiculatum*. Low lying areas are characterised by dense thicket to open Savanna with *Senegalia (Acacia) nigrescens*, *Dichrostachys cinerea*, and *Grewia bicolor* dominating the woody layer, particularly the Granite Lowveld (Mucina and Rutherford, 2006).

Three major land-use types were identified in each of the villages. First, the settlement areas were used for residential purposes (hereafter settlements) (Odhiambo and Magandini, 2008). The majority of households had large gardens (50-80 m x 40-80 m) which were used to grow crops (maize (*Zea mays*), peanuts, beans (*Phaseolus vulgaris*), ground nuts (*Arachis hypogaea*), avocados mangoes, bananas, litchis, and oranges), and to overnight livestock (cattle, donkeys, sheep, goats, and poultry). The second land-use type identified was cropping areas (hereafter crops). Residents of both villages practiced either rotational cropping (maize, ground nuts, and beans) or intercropping (maize, beans, and pumpkins (*Cucurbita* spp.)). Land preparation was usually by manual labour, and preparation typically began in October or November, while planting commenced in early December. Harvesting of crops occurs in February until late April (crop dependant). Farmers reported yields varying between 5 to 20 bags (each bag weighing 50 kg) of maize and 3 to 10 bags of ground nuts (Swanepoel, unpublished data). Crop residues were typically used for livestock fodder. The third land-use type was the grazing areas, which comprised of short grass, shrubs and tall trees (hereafter grazing). In addition to communal grazing of livestock, these areas served for firewood collection and informal hunting. Due to poor land management practices, however, the grazing areas were typically severely overgrazed, with woody plants (mainly *Dichrostachys cinerea*) decreasing herbaceous production and replacing the grass and shrub layer, typically in low lying areas.

2.2. Potential small carnivore diversity and ecosystem services

We define predation of rodent pests and consumption of carrion as potential ecosystem services (Ćirović *et al.*, 2016) that could be provided by small carnivores. We estimated theoretical small carnivore diversity for our study sites by compiling a list of all small carnivore species

potentially present at the study sites from the IUCN Red List of Threatened Species (IUCN, 2016) and from published literature (Apps, 2012; Cillié, 2013; Kingdon and Hoffman, 2012; Skinner and Chimimba, 2005; Stuart and Stuart, 2007). For each species we then extracted from the literature, data on the amount of rodents in their diets, and whether the species consumed carrion (Admasu *et al.*, 2004a, b; Apps, 2012; Camps, 2008; Cillié, 2013; Kingdon and Hoffman, 2012; Skinner and Chimimba, 2005). We regarded species with diets that included a minimum of 20% rodents as potential ecosystem service providers (Ćirović *et al.*, 2016). The home range size of the species potentially present, were used to determine the average distance between camera traps.

2.3. Camera trapping and data preparation

We used camera trapping to determine both species richness and habitat use (occupancy) of small carnivores. Our surveys were underpinned by an occupancy based modelling framework, which guided the layout of camera traps (MacKenzie and Bailey, 2004). Each study area was divided into a settlement area, cropping area and grazing area, based on recent satellite imagery (Google, 2014), which was then overlaid with a regular spaced grid with a cell size of 300 x 300 m (9 ha). The size choice of the grid cells was guided by the median home range size of small carnivores expected to inhabit the study areas (Table 1), to adhere to the independent assumptions of occupancy models (Mackenzie and Royle, 2005). We deployed one camera trap in each grid, which resulted in an average spacing between camera traps of 193 m (standard deviation 65 m), and camera traps were operated for 10-12 days. Camera traps were set to record 24 hours per day, with a 30 second delay between detections. We regarded individuals of the same species photographed within a 5-minute period as the same individual, to avoid pseudo-autocorrelation.

We deployed camera traps at roads, drainage lines, and well-established animal paths. We placed cameras around 30 cm above the ground, and cleared vegetation in front of camera traps to reduce the number of false triggers. In the settlement grid cells we deployed 27-30 infra-red flash cameras (Cuddeback Ambush 1194), as these were less disruptive to the inhabitants of

villages than cameras using a visible light flash, while in the crops and grazing areas we deployed 55-60 xenon flash cameras (Cuddeback Ambush 1170). Camera traps were deployed between 2-26 June 2014 at Ka-Ndengeza and 17 June to 27 July 2014 at Vyeboom. This resulted in a camera trapping effort of 810 trap days in Ka-Ndengeza and 738 trap days in Vyeboom. From each camera trap we extracted detection-non-detection data for the target species, and calculated the relative abundance index (RAI) (O'Brien *et al.*, 2003) of other species we deemed important to the detection and occupancy of target species, such as domestic cats and dogs, livestock, and humans.

To classify land use we first digitized the different land-use types using satellite imagery from Google Maps (Google, 2014), which we later ground-truthed. This approach allowed us to plan the locations of our camera traps for optimal spacing, stratified by land use. We classified crops as either active fields, i.e. still showing agricultural activity, or as abandoned fields. For each camera trap we calculated the percentage of crops, grazing and settlement that comprised the camera trapping grid cell in which each camera trap was located. Camera trap images were catalogued using Camera Base version 1.7 (Tobler, 2015).

2.4. Questionnaires

We assessed the opinions of community members towards small carnivores using a structured questionnaire (Appendix S2) (based on the questionnaire used by Holmern and Røskoft (2014)), completed by a total of 127 respondents (n = 58 in Ka-Ndengeza and n = 69 in Vyeboom). For each camera trap the inhabitants of the nearest household were sampled, but when this was not possible another nearby house was selected. Photographs of small carnivore species were provided to ensure that the species were correctly identified. We asked interviewees whether they had seen each species of carnivore, if they were good for the community, if they kill rodents, if they had impacted the respondents negatively, and if they were aware if any small carnivore species that are killed by people. The reasons for any positive and negative impacts of the species were also recorded. We also asked whether interviewees consider poultry to be an important source of protein, in order to gain some insight into the

motivations for farming chickens and protecting them by killing carnivores.

Ethical approval for the study was provided by the Ethics Committee of the University of Venda (approval number SMNS/14/ZOO/03/2803). We also obtained consent to interview community members of Ka-Ndengeza and Vyeboom from each community Chief in addition to community members. We informed each respondent that anonymity would be maintained, and obtained written consent from interviewees.

2.5. Data analysis

2.5.1. Community occupancy (probability of use) model

We used the MaoTau function in the EstimateS package (Colwell, 2016) to generate species accumulation curves to confirm sampling adequacy for the camera trap dataset (Gotelli and Colwell, 2011). We also used the camera trap data to estimate how the relative abundance of domestic animals influenced small carnivore occupancy, which can be defined as the proportion of the study site that was occupied by the study species (MacKenzie *et al.*, 2017). This is of interest because domestic animals could outcompete sympatric wild carnivores (Vanak and Gompper, 2009), reducing their capacity to provide ecosystem services. Due to the fact that little is known regarding home range and movement rates of South African small carnivores (Roemer *et al.*, 2009), we considered among-grid cell movement in small carnivore species a plausible violation of the closure assumption. As such the occupancy parameter (ψ) should be considered to represent the proportion of area used rather than the proportion of area occupied (MacKenzie and Bailey, 2004).

We adopted the hierarchical formulation of the Dorazio/Royle community occupancy model with data augmentation to estimate species-specific occupancy and site-specific species richness (Dorazio and Andrew Royle, 2005). In a single-species single-season occupancy model the probability that site j is occupied by species z_j is a Bernoulli random variable

governed by the occupancy probability ψ . The occupancy probability is modelled on the logit scale as either a function of site specific covariates or being constant. Analogous to occupancy, the probability that a species is detected is governed by the detection probability, p , which is conditioned on the true latent occupancy state, z_j . Survey sites are camera trapped on k occasions (e.g. days) where the observations, y_{jk} , is a Bernoulli random variable, either $p_{jk} = 1$ where $z_j = 1$ or $p_{jk} = 0$ where $z_j = 0$. Detection probability is also modelled on the logit scale, either constant or as a function of site (e.g. vegetation type) or occasion (e.g. daily temperature) specific covariates.

We fitted community models to the data, as this allowed us to investigate the influence of the relative abundance of domestic animals on small carnivores at a community level (MacKenzie *et al.*, 2017). In the community model formulation the single-species single-season model is further extended where the latent and model parameters are indexed by species, i . This formulation results in a number of linked species-specific models because it is assumed that these species-specific parameters come from a common underlying distribution (governed by the hyperparameters, which in our study is the small carnivore community). To estimate the number species at each sampling site (including ones never detected) we augmented the data with all-zero observations for the hypothetical species (Dorazio and Andrew Royle, 2005). We hypothesized that in our study area a potential 23 small carnivore species could occur (IUCN, 2016), and we therefore augmented the observed data with 14 species.

We expected occupancy and diversity of small carnivores to be affected by various anthropogenic and environmental variables. To investigate these variables we developed an *a priori* model based on biological hypotheses on how small carnivore occupancy could be influenced by these variables. We hypothesized that small carnivore occupancy will be affected by the presence of domestic cat, dogs, livestock, humans and land use. Both domestic cats and dogs can either directly (through predation) or indirectly (through competitive exclusion) impact small carnivores (Brook *et al.*, 2012; Dickman, 1996). Similarly, humans can directly kill small carnivores (Berger, 2006; Ćirović *et al.*, 2016), and livestock can trample burrows of

small carnivores and reduce vegetation cover (Blaum *et al.*, 2007a; Blaum *et al.*, 2007b). We used variance inflation factor (Zuur *et al.*, 2009) to identify and remove highly correlated variables to reduce multicollinearity. Using all the covariates we sequentially dropped the variable with highest VIF (however, we selected the variable with the least biological effect among variables with high VIF first), and recalculated the VIF until the VIF of each factor was below five (Zuur *et al.*, 2009). Using this approach we dropped percentage crops, settlement and grazing as these variables were highly correlated and had high VIF factors. Both human RAI and dog RAI were correlated and we thus dropped human RAI since we hypothesised that domestic dogs can have higher sustained impact on small carnivores (e.g. since dogs can roam over the landscape independent of humans).

We thus retained only domestic cat RAI, domestic dog RAI, and livestock RAI as explanatory occupancy covariates, and we modelled occupancy probability as having species-specific random intercepts with these three site covariates. We assumed that occupancy patterns were similar across villages, even though they were not sampled at the same time. For detection probability we only modelled the effect of survey date (Julian day) on detection, again as species-specific random intercept (Dorazio and Andrew Royle, 2005). We collapsed the 10-12 day survey into 5 sampling occasions to increase detection probabilities (Ramesh *et al.*, 2012), and each camera trap was regarded as independent.

We used a Bayesian framework (Plummer, 2003) to implement the community model. Full details can be found in Appendix S3, while the full model specification can be found in Appendix S4. Results are reported in mean, standard deviation and 95% Bayesian confidence intervals (95 BCI taken from the 2.5% and 97.5% percentiles of the posterior mean). We regarded coefficients as having strong inference value if its 95 BCI values did not include 0. We further estimated the number of small carnivore species per land use by summing the estimated species richness at each survey site, in each land use. Finally we used the estimated species richness at each camera trap location to create spatially explicit species richness maps using inverse distance weighted interpolation (Sarmiento *et al.*, 2010). We used R v3.4.1 (R

Development Core Team, 2017) for all modelling, with the following R packages; raster for IWD (Hijmans, 2015), jagsUI (Kellner, 2016).

2.5.2. Questionnaires

The questionnaire data allowed us to investigate stakeholder perceptions of small carnivores in agro-ecosystems. We explored the questionnaire data by calculating the frequency with which respondents reported that 1) they had seen small carnivores; 2) small carnivores had either positive or negative impacts on people; 3) small carnivores kill rodents; and 4) people kill small carnivores. Some frequencies were represented graphically using bar plots created using the R package ggplot2 (Wickham, 2016). All data analysed in this study are publically available in Williams *et al.* (2017).

3. Results

3.1. Small and medium carnivore diversity and occupancy (probability of use)

Species accumulation curves plateaued at approximately 1,368 camera trapping days (8 survey days), which suggested adequate sampling (Appendix S5). Of 23 small and medium carnivore species potentially occurring at the study sites (IUCN, 2016), we detected 9 (8 at Ka-Ndengeza and 8 at Vyeboom) small carnivores representing 5 different families (Table 1). The mean metacommunity richness was estimated at 14.48 (95 BCI 9-22 species). However the mean metacommunity richness had a skewed posterior distribution and a wide credible interval. We therefore used the mode to estimate total metacommunity richness, which was estimated at 10.98 species.

The strength of associations with occupancy covariates varied between species (Fig. 1). The presence of cats did not have a strong association with any of the small and medium carnivore species, nor to the metacommunity as a whole (Fig. 1). In contrast, dogs had a strong negative

association with occupancy probability (probability of use) for all species and the metacommunity (Fig. 1). For livestock only four species (white tailed mongoose, slender mongoose, Selous' mongoose, and large spotted genet) showed strong positive associations with livestock presence, while the other five species had no association. Interestingly, the metacommunity also had a strong positive association with livestock presence (Fig. 1).

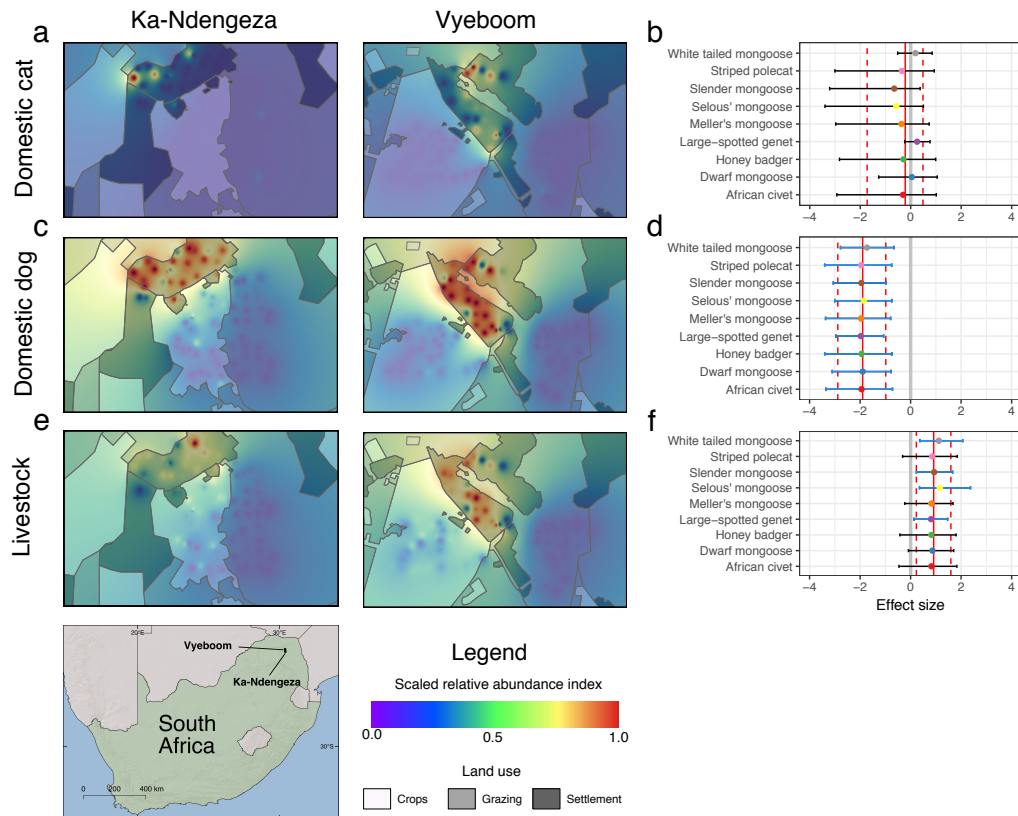


Fig. 1. Interpolated heat maps based on relative abundance index (scaled between 0 and 1) for a) domestic cat, c) domestic dog, and e) livestock across the settlement, crop, and grazing areas in Ka-Ndengeza and Vyeboom. Caterpillar plots show the strength of associations between the RAI of b) domestic cat, d) domestic dog, and f) livestock with occupancy (probability of use) of the nine carnivore species detected. Confidence intervals highlighted in blue do not overlap 0. The broken lines indicate the 95% BCI for the mean community response to each variable.

Cropping areas consistently showed higher species richness than grazing and settlement areas (Fig. 2). Spatially, species richness density surfaces clearly adhered to cropping areas and highest species richness per 900 m² grid cell were consistently observed in the cropping areas (Fig. 2). A survey of the literature showed that 65% of these species (15/23) are reported to have at least 20% of rodents in their diet (Table 1). Combined with species richness maps this suggests that the small and carnivore community not only occur most often in cropping areas, but also probably incorporate a large proportion of rodents in their diet. Using the mode small carnivore richness (10.98) as a reliable estimate of species richness we suggest that the study area realised around 47% of the potential small carnivore diversity.

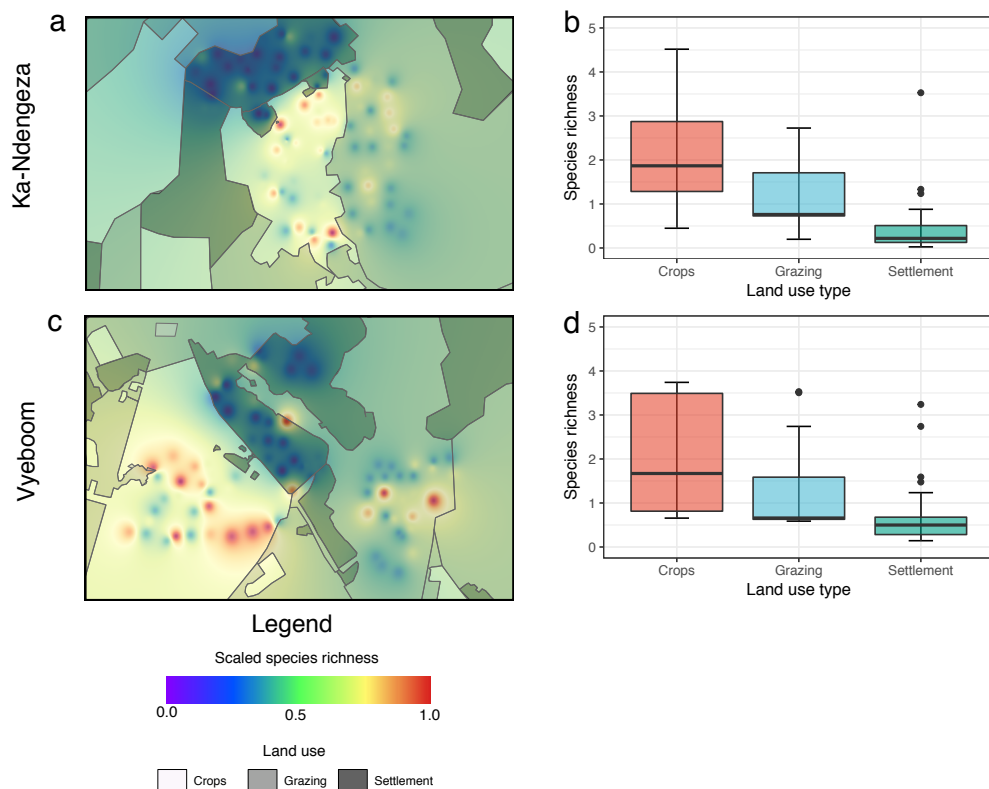


Fig. 2. Maps and boxplots showing how the species richness (scaled between 0 and 1) of small carnivores varies with land use at Ka-Ndengeza (a, b) and Vyeboom (c, d). Boxplots show mean number (posterior mean) of species estimated at each camera trap, summarized per land use.

337 Table 1. List of carnivore species detected during the camera trap study. The table is ordered according to family level (all capitals).

					Number of independent detections per 1,000 camera trap days						
Common name	Scientific name	Home range size (km²)	Consumes carrion	% of scats or stomachs that contain rodent remains	Ka-Ndengeza			Vyeboom			IUCN Red List ⁵
					Settlement	Crops	Grazing	Settlement	Crops	Grazing	
CANIDAE	Domestic dog <i>Canis lupus familiaris</i>				9324.1	1269.8	308.1	5160	201.7	37.04	
MUSTELIDAE											
Striped polecat	<i>Ictonyx striatus</i>	-	No	20-30 ¹	0	0	5.1	0	8.23	0	Least concern
Honey badger	<i>Mellivora capensis</i>	10 - 30	Yes	30 ¹ , 57 ²	0	0	0	0	0	6.17	Least concern
FELIDAE											
Domestic cat	<i>Felis catus</i>				324.07	0	10.1	720	0	6.14	
VIVERRIDAE											

Large-spotted genet	<i>Genetta maculata</i>	0.5 - 1	No	47 ³ , 68 ⁴	0	642.86	217.17	22.22	172.8	228.4	Least concern
African civet	<i>Civettictis civetta</i>	5 - 11.1	Yes	41 ⁴	0	0	0	0	8.23	0	Least concern
HERPESTIDAE											
Slender mongoose	<i>Galerella sanguinea</i>	0.5 - 1	Yes	25 ³	0	253.97	25.25	0	148.15	86.42	Least concern
Meller's mongoose	<i>Rhynchogale melleri</i>	-	No	Not available	0	47.62	0	0	0	0	Least concern
Selous' mongoose	<i>Paracynictis selousi</i>	-	No	Not available	0	71.43	0	0	32.92	0	Least concern
White tailed mongoose	<i>Ichneumia albicauda</i>	4 - 8	Yes	18 ³	0	150.79	0	26.67	8.23	18.52	Least concern
Dwarf mongoose	<i>Helogale parvula</i>	1 - 3	No	4	0	31.75	0	4.44	4.12	30.86	Least concern
Species richness					2	7	5	5	8	7	
% of potential maximum species richness (23)					9	30	22	22	35	30	

338

339

340

341 ¹Apps (2012)

342 ²Skinner and Chimimba (2005)

343 ³Smithers (1971)

344 ⁴Smithers and Wilson (1979)

345 ⁵IUCN (2016)

346

347 3.2. Questionnaires

348 Eleven species of non-domesticated small carnivore species were reported to be seen by the
349 respondents (Appendix S6). All mongoose species (with the exception of water mongoose),
350 African wildcat, small spotted genet, black backed jackal, and striped polecat were reported
351 most frequently. African civet and honey badger were seen by few respondents, while caracal,
352 serval, and water mongoose had not been seen. Domestic cats and domestic dogs had been seen
353 by all interviewees. The only species perceived to benefit the community were domestic cats
354 and domestic dogs (Table 2).

355 Table 2. Percentage of respondents (n = 58 in Ka-Ndengeza and n = 69 in Vyeboom) with positive responses to questions on interactions between
 356 carnivores and humans.

Species	Are they good for the community?		Do they kill rodents?		Do they impact you negatively?		Do people kill them?	
	Ka-Ndengeza	Vyeboom	Ka-Ndengeza	Vyeboom	Ka-Ndengeza	Vyeboom	Ka-Ndengeza	Vyeboom
Banded mongoose	0	0	0	15.9	20.7	43.5	0	0
Dwarf mongoose	0	0	5.2	15.9	32.8	95.7	1.7	1.4
Slender mongoose	0	0	25.9	15.9	89.7	79.7	8.6	0
Yellow mongoose	0	0	1.7	11.6	0	0	1.7	0
White tailed mongoose	0	0	3.4	15.9	22.4	72.5	0	0
Water mongoose	0	0	0	0	0	0	0	0
Black backed jackal	0	0	0	0	0	5.8	0	0
African civet	0	0	0	0	0	0	0	0
Small spotted genet	0	0	13.8	0	1.7	0	0	0
Striped polecat	0	0	27.6	0	0	0	0	0
Caracal	0	0	0	0	0	0	0	0
African wild cat	0	0	44.8	62.3	6.9	43.5	1.7	0
Honey badger	0	0	0	0	0	0	0	0
Domestic cat	51.7	98.6	100	100	6.9	1.4	0	0
Domestic dog	58.6	98.6	3.4	0	8.6	1.4	0	0

357

A total of eight species of non-domesticated carnivores were believed by some people to kill rodents (Ka-Ndengeza: seven species were thought to kill rodents by a mean of 17.5% of respondents; Vyeboom: six species were thought to kill rodents by a mean of 23.0% of respondents). The species most commonly thought to predate on rodents were African wildcat, striped polecat, and slender mongoose (Table 2).

Negative impacts of carnivores on people were reported for most mongoose species, black backed jackal, small spotted genet, and African wild cat (Table 2). Most negative impacts were perceived to be due to poultry predation, although a small number of respondents cited cultural reasons, such as involvement in witchcraft or other superstitions, for negative impacts (Appendix S7).

Slender mongoose, dwarf mongoose, yellow mongoose, and African wildcat were said to be killed by people (Table 2). The only reason provided for people killing carnivores was poultry predation. Poultry was considered to be an important source of protein by 98.3% of respondents in Ka-Ndengeza and 100.0% of respondents in Vyeboom. The median number of chickens owned was 10 (interquartile range = 13, n = 21) in Ka-Ndengeza, and 4 (interquartile range = 6, n = 24) in Vyeboom. Poultry were almost always free-ranging (in 96.6% and 100% of households surveyed in Ka-Ndengeza and Vyeboom respectively).

4. Discussion

Our camera trapping results indicated that cropping areas consistently supported the greatest diversity of small carnivores. Furthermore, the literature review showed that the small carnivore assemblages present typically incorporate a large percentage of rodents and carrion in their diets. Collectively these results highlight the potential for pest control and carrion removal by small carnivores as important ecosystem services. Our results concur with other studies that highlight the unrealised potential of small carnivore predation and scavenging as ecosystem services (Ćirović *et al.*, 2016; Mateo-Tomás *et al.*, 2015). Rodent pests, for example,

account for approximately 15% of the damage caused to rural farming crops in Africa (Swanepoel *et al.*, 2017), and such damage is dependent on the density of rodents (Brown *et al.*, 2007). Since small carnivore diets include a large proportion of rodents, it is likely that small carnivore predation could be a key factor affecting rodent abundance, and therefore reduce crop damage (Ćirović *et al.*, 2016). Further support comes from meta-analysis studies, that show that reduced predation increases population growth for cyclic prey (Salo *et al.*, 2010) and provisioned populations of small mammals such as rodents feeding on grain (Prevedello *et al.*, 2013; Salo *et al.*, 2010). There therefore appears to be strong support, both from our findings and from the literature, that predation of rodents by small carnivores could be an important ecosystem service to rural communities through EBRM.

Our results showed that abundance of domestic dogs (and feral dogs) and livestock are important determinants of small carnivore diversity and habitat use, while cats seemed to have little effect. Several studies have highlighted the negative impact of dogs (domestic and feral) on native mammalian communities (Hughes and Macdonald, 2013; Reed and Merenlender, 2011). For example, dogs can act as intraguild competitors where they can outcompete carnivores, especially under conditions of low prey biomass (Vanak and Gompper, 2009). We suggest that such a scenario is most likely prevalent in rural African landscapes where local fauna often form part of the diet of people in rural areas (Holmern *et al.*, 2006). Furthermore dogs, especially when roaming freely (a scenario common in African rural landscapes (Czupryna *et al.*, 2016)), can kill small carnivores (Ralls and White, 1995). Finally, dogs are often used during hunting activities where they can kill non-target species such as small carnivores (Holmern *et al.*, 2006).

The lack of effect of cats on small carnivore occupancy is surprising, given the large impact cats have on mammalian communities (Loss *et al.*, 2013). We provide two possible reasons for this lack of effect; first cats most often include small mammals in their diet (Loss *et al.*, 2013), and as such might impact small carnivores through competitive exclusion (Brook *et al.*, 2012). However, densities of cats in our study might not be high enough to achieve such an effect.

Secondly, dog hunting often occurs at night (Holmern *et al.*, 2006), which might restrict cats (and hence their impact on small carnivores) to the settlement areas. The positive effect of livestock contrasts with other studies that highlight the negative impact of livestock on small carnivores (Blaum *et al.*, 2007a; Blaum *et al.*, 2007b). We hypothesised that this effect is probably mediated through invertebrate food sources for small carnivores. For example the four small carnivore species exhibiting a positive occupancy effect due to livestock (large spotted genet, slender mongoose, white tailed mongoose and Selous' mongoose) all incorporate a large proportion of invertebrates in their diet (Skinner and Chimimba, 2005). Studies have shown that disturbance-adapted insect populations increase in abundance in highly impacted areas (e.g. heavy grazed) (Schowalter, 1985; Seymour and Dean, 1999). Therefore, the presence of livestock can create local conditions of increased invertebrate biomass, which could facilitate small carnivore presence.

We found that cropping areas had the highest small carnivore richness, which contrasts with the low biodiversity often observed in intensive agricultural systems (Benton *et al.*, 2003). We provide several hypotheses for this observation, which are not necessarily mutually exclusive. First, rural agricultural landscapes are often structurally complex and heterogeneous (Donald, 2004) which seems to support higher animal diversity (Norris, 2008). Secondly, rural agricultural systems support a diverse and high rodent abundance, especially in our study areas (Belmain, 2006), which can support small carnivores (Blaum *et al.*, 2007b). While dogs had a large effect on small carnivores, the highest dog and cat activities were observed in the settlement areas, and to a lesser extent in the cropping areas, which suggests that competitive exclusion and competition with small carnivores (Glen and Dickman, 2005; Vanak and Gompper, 2010) is limited in agricultural areas. Finally livestock abundance was higher in cropping areas compared to grazing areas, which could have created favourable conditions for high biomass of disturbance-adapted insect populations that can act food resource for small carnivores (Seymour and Dean, 1999).

While our results support the hypothesis that small carnivores could provide ecosystem

services, we highlight that such a service would not depend solely on diversity, but also abundance of small carnivores. Our results show that the majority of small carnivores had low relative abundance indices, which were likely to be below ecologically effective densities (Soulé *et al.*, 2005). Nonetheless, the small carnivore assemblage present in these rural agro-ecosystems can still fulfil basic ecological functionality of predation (Roemer *et al.*, 2009). Such functionality will be largely dependent on whether the small carnivore assemblages retained inherent functional redundancy (Roemer *et al.*, 2009; Suraci *et al.*, 2017). This is important since the ecosystem service provision can be greater if expressed through collective effects, where the sum effect of predation (from different carnivores) might exceed that of a single small carnivore (Suraci *et al.*, 2017). Our study shows that the system retained some functional redundancy, however a large number of rodent specialists (e.g. striped polecat) were not detected or occurred at low relative abundances. Their absence probably reflects the small carnivore assemblage responding to pressures and changes as a result of human modification to the landscape that exist around rural agro-ecosystems. These responses will inadvertently bring shifts and changes in ecosystem service delivery and provision, which, if not checked can ultimately only exist as simple linear food chain communities (Roemer *et al.*, 2009). Therefore facilitating or at least maintaining small carnivore functional redundancy should be a key conservation management action in rural African landscapes if ecosystem services are to be maintained. Changes in rural landscapes are dynamic, which could potentially allow for various species of small carnivores to persist in them (Melo *et al.*, 2013). However, to what extent these changes retain or enhance functional redundancy remains to be explored.

Encouragingly, community members were able to identify 11 native small carnivore species that should occur in their areas, although we recorded fewer species using camera traps (nine wild species, domestic cats and domestic dogs). Although respondents were aware of the presence of the study species in their villages, and many respondents acknowledged the presence of rodents in the diet of some wild small carnivore species, they lacked any appreciation of the ecosystem services that they could provide. Reports of negative impacts of small carnivores were commonplace, almost exclusively due to perceived poultry predation. In both villages keeping of poultry was very common, and almost all respondents asserted that

poultry was an important source of protein in their diet. The threat of poultry predation was said to be the main motivation for small carnivores being killed by community members.

The mechanism by which some small carnivores were thought to predate on poultry was unconventional and unsubstantiated. Many community members believed that carnivores would intentionally trap the beaks of chickens in their anus, before breaking their necks. Although some species of small carnivores such as the African civet, small spotted genet, and large spotted genet have been known to predate on poultry (Kingdon and Hoffman, 2012), and in some cases levels of poultry predation by small carnivores can be high (Holmern and Røskoft, 2014), such perceptions illustrate that the perceived threats of predation may not always have a strong grounding in reality. Nevertheless, it appears that overcoming perceptions of poultry predation will be the key challenge in promoting the role of small carnivores as providers of ecosystem services. Our results could help to demonstrate to community members that wild small carnivores are more likely benefit them by controlling pests and removing carcasses than predate on their poultry. We note that the wording of the questionnaires (Holmern and Røskoft, 2014) could be improved upon to reduce bias. As an example, we suggest that in future studies asking respondents to rate their benefit of a carnivore species on a Likert scale would be less biased than asking if a species is good for the community (Morgan-Brown *et al.*, 2010).

Although our findings indicate that small carnivores could provide ecosystem services through pest control and waste removal in rural agro-ecosystems, we suggest that further research may help to characterise the impacts of small carnivores on the density and diversity of rodents in agricultural fields, the amount of crop damage caused by rodents, and the amount of carrion removed. The socio-economic implications on the livelihoods of people adopting these strategies would also be worthy of further study.

5. Conclusions

Our findings suggest that agricultural areas could be important refuges for small carnivores within modified landscapes, and these species are likely to be providing important ecosystem services in rural agro-ecosystems. We found that agricultural areas supported the greatest diversity of small carnivores. Livestock was linked to higher levels of occupancy (probability of use) of small carnivores, while the opposite trend was observed for domestic dogs, and domestic cats had no influence on carnivore occupancy. The small carnivore species present are reported in the literature to dedicate a considerable proportion of their diets to rodents, and consume carrion. Although community members could identify many small carnivore species, they appeared to be unaware of the ecosystem services that the small carnivores are likely to provide through EBRM and carcass removal. The perceived threat of poultry predation emerged as a key challenge in promoting the role of small carnivores as providers of ecosystem services.

6. Appendices

Appendix S1. Study area figure

Appendix S2. Interview schedule.

Appendix S3. Model description and parameter estimates of the community occupancy model applied to small carnivore camera trapping data from a rural matrix.

Appendix S4. Community model JAGS code used in the analysis.

Appendix S5. Species accumulation curves to show sampling adequacy.

Appendix S6. Percentage of respondents in Ka-Ndengeza and Vyeboom that reported seeing species of small carnivores.

Appendix S7. Reasons provided why carnivores have impacted respondents negatively for Ka-Ndengeza and Vyeboom.

7. Acknowledgements

We are grateful to the residents of Ka-Ndengeza and Vyeboom for granting permission to collect these data, and for their support and hospitality. We are particularly grateful to the participants that completed the questionnaires, and the residents who allowed us to place camera traps on their land. The research has primarily been funded by the Sasol Agricultural Trust (South Africa), Univen Niche fund (SMNS /17/Zoo/01), International Foundation for Science (D/4984-2), the European Union through its ACP S & T programme (StopRats; FED2013-330223; <http://www.acp-hestre.eu/>) with further funding from the German Federal Ministry of Education and Science (“BMBF”) through the project “Limpopo Living Landscapes - Understanding the Dynamics of Ecological and Cultural Landscapes, in the Face of Global Change, in the Northern Limpopo Region of South Africa” of the SPACES (Science Partnerships for the Assessment of Complex Earth Systems) consortium (https://www.fona.de/mediathek/pdf/SPACES_Broschuere_Englisch.pdf). PJT acknowledges the support of the University of Venda, the National Research Foundation, and the Department of Science and Technology under the South African Research Chairs Initiative (SARChI) on Biodiversity Value and Change within the Vhembe Biosphere Reserve, hosted at University of Venda and co-hosted by the Centre for Invasion Biology at University of Stellenbosch. We thank Matt Hayward and one anonymous reviewer for their comments and suggestions which improved the manuscript. We also thank Mike Meridith.

8. References

- Admasu, E., Thirgood, S.J., Bekele, A. and Laurenson, M.K., 2004a. A note on the spatial ecology of African civet *Civettictis civetta* and common genet *Genetta genetta* in farmland in the Ethiopian Highlands. *Afr. J. Ecol.* **42**, 160-162.
- Admasu, E., Thirgood, S.J., Bekele, A. and Laurenson, M.K., 2004b. Spatial ecology of white-tailed mongoose in farmland in the Ethiopian Highlands. *Afr. J. Ecol.* **42**, 153-159.
- Apps, P., 2012. *Smithers' Mammals of Southern Africa: A Field Guide*. Random House Struik, Cape Town.

558

559 Belmain, S.R., 2006. Rats and human health in Africa: proceedings of an international
560 workshop on rodent-borne diseases and the RatZooMan research project. The Natural
561 Resources Institute, Greenwich.

562

563 Benton, T.G., Vickery, J.A. and Wilson, J.D., 2003. Farmland biodiversity: is habitat
564 heterogeneity the key? *Trends Ecol. Evol.* **18**, 182-188.

565

566 Berger, K.M., 2006. Carnivore-livestock conflicts: effects of subsidized predator control and
567 economic correlates on the sheep industry. *Conserv. Biol.* **20**, 751-761.

568

569 Blaum, N., Rossmanith, E., Popp, A. and Jeltsch, F., 2007a. Shrub encroachment affects
570 mammalian carnivore abundance and species richness in semiarid rangelands. *Acta Oecol.* **31**,
571 86-92.

572

573 Blaum, N., Rossmanith, E., Schwager, M. and Jeltsch, F., 2007b. Responses of mammalian
574 carnivores to land use in arid savanna rangelands. *Basic Appl. Ecol.* **8**, 552-564.

575

576 Blaum, N., Tietjen, B. and Rossmanith, E., 2009. Impact of livestock husbandry on small-and
577 medium-sized carnivores in Kalahari savannah rangelands. *J. Wildlife Manage.* **73**, 60-67.

578

579 Brook, L.A., Johnson, C.N. and Ritchie, E.G., 2012. Effects of predator control on behaviour
580 of an apex predator and indirect consequences for mesopredator suppression. *J. Appl. Ecol.* **49**,
581 1278-1286.

582

583 Brown, P.R., Huth, N.I., Banks, P.B. and Singleton, G.R., 2007. Relationship between
584 abundance of rodents and damage to agricultural crops. *Agric. Ecosyst. Environ.* **120**, 405-415.

585

586 Brown, P.R., Tuan, N.P., Singleton, G.R., Ha, P.T.T., Hoa, P.T., Hue, D.T., Tan, T.Q., Van
587 Tuat, N., Jacob, J. and Müller, W.J., 2006. Ecologically based management of rodents in the
588 real world: Applied to a mixed agroecosystem in Vietnam. *Ecol. Appl.* **16**, 2000-2010.

589

590 Buckle, A.P. and Smith, R.H., 2015. Rodent pests and their control. CABI, Wallingford.

591

592 Camps, D., 2008. Activity patterns of adult common genets *Genetta genetta* (Linnaeus, 1758)
593 in northeastern Spain. *Galemys* **20**, 47-60.

594

595 Cancio, I., González-Robles, A., Bastida, J.M., Isla, J., Manzaneda, A.J., Salido, T. and Rey,
596 P.J., 2017. Landscape degradation affects red fox (*Vulpes vulpes*) diet and its ecosystem
597 services in the threatened *Ziziphus lotus* scrubland habitats of semiarid Spain. *J. Arid Environ.*
598 **145**, 24-34.

599

600 Caro, T. and Stoner, C., 2003. The potential for interspecific competition among African
601 carnivores. *Biol. Conserv.* **110**, 67-75.

602

603 Cillié, B., 2013. *The Mammal Guide of Southern Africa*. Briza, Pretoria.

604

605 Ćirović, D., Penezić, A. and Krofel, M., 2016. Jackals as cleaners: Ecosystem services provided
606 by a mesocarnivore in human-dominated landscapes. *Biol. Conserv.* **199**, 51-55.

607

608 Colwell, R.K., 2016. EstimateS: Statistical estimation of species richness and shared species
609 from samples. v9.1.0 [computer program]. Available from <http://purl.oclc.org/estimates>.

610

611 Crooks, K.R., 2002. Relative sensitivities of mammalian carnivores to habitat fragmentation.
612 *Conserv. Biol.* **16**, 488-502.

613

614 Czapryna, A.M., Brown, J.S., Bigambo, M.A., Whelan, C.J., Mehta, S.D., Santymire, R.M.,
615 Lankester, F.J. and Faust, L.J., 2016. Ecology and demography of free-roaming domestic dogs
616 in rural villages near Serengeti National Park in Tanzania. *PLoS One* **11**, e0167092.

617

618 Dickman, C.R., 1996. Overview of the impacts of feral cats on Australian native fauna.
619 Australian Nature Conservation Agency, Canberra.

620

621 Dobson, A., Lodge, D., Alder, J., Cumming, G.S., Keymer, J., McGlade, J., Mooney, H.,
622 Rusak, J.A., Sala, O. and Wolters, V., 2006. Habitat loss, trophic collapse, and the decline of
623 ecosystem services. *Ecology* **87**, 1915-1924.

624

625 Donald, P.F., 2004. Biodiversity impacts of some agricultural commodity production systems.
626 *Conserv. Biol.* **18**, 17-38.

627

628 Dorazio, R.M. and Andrew Royle, J., 2005. Estimating size and composition of biological
629 communities by modeling the occurrence of species. *J. Am. Stat. Assoc.* **100**, 389-398.

630

631 Glen, A.S. and Dickman, C.R., 2005. Complex interactions among mammalian carnivores in
632 Australia, and their implications for wildlife management. *Biol. Rev. Camb. Philos. Soc.* **80**,
633 387-401.

634

635 Google, 2014. Satellite imagery. Sources: Landsat/Copernicus. Image date 31 December 2013.
636 Available from <https://www.google.co.za/maps>.

637

638 Gotelli, N.J. and Colwell, R.K., 2011. Estimating species richness, In *Biological Diversity:*
639 *Frontiers in measuring biodiversity.* eds. Magurran, A.E. and McGill, B.J., pp. 39-54. Oxford
640 University Press, Oxford.

641

642 Granjon, L. and Duplantier, J.-M., 2009. Les rongeurs de l'Afrique sahélo-soudanienne. IRD,
643 Marseille.

644

645 Gusset, M., Swarner, M., Mponwane, L., Keletile, K. and McNutt, J., 2009. Human–wildlife
646 conflict in northern Botswana: livestock predation by endangered African wild dog *Lycaon*
647 *pictus* and other carnivores. *Oryx* **43**, 67-72.

648

649 Hijmans, R.J., 2015. Geographic data analysis and modelling. R package version 2.5-8
650 [computer program]. Available from <http://cran.univ-lyon1.fr/web/packages/raster/>.

651

652 Hijmans, R.J., Cameron, S.E., Parra, J.L., Jones, P.G. and Jarvis, A., 2005. Very high resolution
653 interpolated climate surfaces for global land areas. *Int. J. Climatol.* **25**, 1965-1978.

654

655 Holmern, T., Mkama, S., Muya, J. and Røskoft, E., 2006. Intraspecific prey choice of bushmeat
656 hunters outside the Serengeti National Park, Tanzania: a preliminary analysis. *Afr. Zool.* **41**,

81-87.

Holmern, T. and Røskoft, E., 2014. The poultry thief: Subsistence farmers' perceptions of depredation outside the Serengeti National Park, Tanzania. *Afr. J. Ecol.* **52** (3), 334-342

Hughes, J. and Macdonald, D.W., 2013. A review of the interactions between free-roaming domestic dogs and wildlife. *Biol. Conserv.* **157**, 341-351.

IUCN, 2016. The IUCN Red List of Threatened Species. Version 2016-3. Available from <http://www.iucnredlist.org/>. Accessed 2016.

Jacob, J., Sudarmaji, Singleton, G.R., Rahmini, Herawati, N.A. and Brown, P.R., 2010. Ecologically based management of rodents in lowland irrigated rice fields in Indonesia. *Wildl. Res.* **37**, 418-427.

Kellner, K., 2016. jagsUI: A wrapper around 'rjags' to streamline 'JAGS' analyses. R package version 1.4.4 [computer program]. Available from <https://cran.r-project.org/package=jagsUI>.

Kingdon, J. and Hoffman, M., 2012. *Mammals of Africa*. Bloomsbury, New York.

Krijger, I.M., Belmain, S.R., Singleton, G.R., Groot Koerkamp, P.W. and Meerburg, B.G., 2017. The need to implement the landscape of fear within rodent pest management strategies. *Pest Manag. Sci.* **73**, 2397–2402.

Labuschagne, L., Swanepoel, L.H., Taylor, P.J., Belmain, S.R. and Keith, M., 2016. Are avian predators effective biological control agents for rodent pest management in agricultural systems? *Biol. Control* **101**, 94-102.

Lamarque, F., Anderson, J., Fergusson, R., Lagrange, M., Osei-Owusu, Y. and Bakker, L., 2009. *Human-wildlife conflict in Africa: causes, consequences and management strategies*. Food and Agriculture Organization of the United Nations, Rome.

689 Loss, S.R., Will, T. and Marra, P.P., 2013. The impact of free-ranging domestic cats on wildlife
690 of the United States. *Nat. Commun.* **4**, 1396.

691

692 Lowder, S.K., Scoet, J. and Raney, T., 2016. The number, size, and distribution of farms,
693 smallholder farms, and family farms worldwide. *World Dev.* **87**, 16-29.

694

695 MacKenzie, D.I. and Bailey, L.L., 2004. Assessing the fit of site-occupancy models. *J. Agric.*
696 *Biol. Environ. Stat.* **9**, 300-318.

697

698 MacKenzie, D.I., Nichols, J.D., Royle, J.A., Pollock, K.P., Bailey, L.L. and Hines, J.E., 2017.
699 *Occupancy Estimation and Modeling: Inferring Patterns and Dynamics of Species Occurrence*,
700 Second edition edn. Academic Press, Oxford.

701

702 Mackenzie, D.I. and Royle, J.A., 2005. Designing occupancy studies: general advice and
703 allocating survey effort. *J. Appl. Ecol.* **42**, 1105-1114.

704

705 Mahlaba, T.a.A.M., Monadjem, A., McCleery, R. and Belmain, S.R., 2017. Domestic cats and
706 dogs create a landscape of fear for pest rodents around rural homesteads. *PLoS One* **12**,
707 e0171593.

708

709 Massawe, A.W., Mulungu, L.S., Makundi, R.H., Dlamini, N., Eiseb, S.J., Kirsten, F., Mahlaba,
710 T., Malebane, P., Von Maltitz, E., Monadjem, A., Taylor, P., Tutjavi, V. and Belmain, S.R.,
711 2011. Spatial and temporal population dynamics of rodents in three geographically different
712 regions in Africa: Implication for ecologically-based rodent management. *Afr. Zool.* **46**, 393-
713 405.

714

715 Masters, W.A., Djurfeldt, A.A., De Haan, C., Hazell, P., Jayne, T., Jirström, M. and Reardon,
716 T., 2013. Urbanization and farm size in Asia and Africa: implications for food security and
717 agricultural research. *Glob. Food Sec.* **2**, 156-165.

718

719 Mateo-Tomás, P., Olea, P.P., Moleón, M., Vicente, J., Botella, F., Selva, N., Viñuela, J. and
720 Sánchez-Zapata, J.A., 2015. From regional to global patterns in vertebrate scavenger
721 communities subsidized by big game hunting. *Divers. Distrib.* **21**, 913-924.

723 Melo, F.P., Arroyo-Rodríguez, V., Fahrig, L., Martínez-Ramos, M. and Tabarelli, M., 2013.
 724 On the hope for biodiversity-friendly tropical landscapes. *Trends Ecol. Evol.* **28**, 462-468.
 725

726 Monadjem, A., Taylor, P.J., Denys, C. and Cotterill, F.P.D., 2015. Rodents of Sub-Saharan
 727 Africa: A biogeographic and taxonomic synthesis. Walter de Gruyter, Berlin.
 728

729 Morgan-Brown, T., Jacobson, S.K., Wald, K. and Child, B., 2010. Quantitative assessment of
 730 a Tanzanian integrated conservation and development project involving butterfly farming.
 731 *Conserv Biol* **24**, 563-572.
 732

733 Mucina, L. and Rutherford, M.C., 2006. The vegetation of South Africa, Lesotho and
 734 Swaziland. South African National Biodiversity Institute, Pretoria.
 735

736 Norris, K., 2008. Agriculture and biodiversity conservation: opportunity knocks. *Conserv. Lett*
 737 **1**, 2-11.
 738

739 O'Brien, T.G., Kinnaird, M.F. and Wibisono, H.T., 2003. Crouching tigers, hidden prey:
 740 Sumatran tiger and prey populations in a tropical forest landscape. *Anim. Conserv.* **6**, 131-139.
 741

742 Odhiambo, J.J.O. and Magandini, V.N., 2008. An assessment of the use of mineral and organic
 743 fertilizers by smallholder farmers in Vhembe district, Limpopo province, South Africa. *Afr. J.*
 744 *Agr. Res.* **053**, 357-362.
 745

746 Plummer, M., 2003. JAGS: A program for analysis of Bayesian graphical models using Gibbs
 747 sampling, In *Proceedings of the 3rd international workshop on distributed statistical*
 748 *computing*. Vienna.
 749

750 Prevedello, J.A., Dickman, C.R., Vieira, M.V. and Vieira, E.M., 2013. Population responses
 751 of small mammals to food supply and predators: a global meta-analysis. *J. Anim. Ecol.* **82**,
 752 927-936.
 753

754 Prugh, L.R., Stoner, C.J., Epps, C.W., Bean, W.T., Ripple, W.J., Laliberte, A.S. and Brashares,
 755 J.S., 2009. The rise of the mesopredator. *Bioscience* **59**, 779-791.

756

757 R Development Core Team, 2017. R: a language and environment for statistical computing
758 [computer program]. Available from <https://cran.r-project.org/>.

759

760 Ralls, K. and White, P.J., 1995. Predation on San Joaquin kit foxes by larger canids. J.
761 Mammal. **76**, 723-729.

762

763 Ramesh, T., Kalle, R., Sankar, K. and Qureshi, Q., 2012. Dry season factors determining
764 habitat use and distribution of mouse deer (*Moschiola indica*) in the Western Ghats. Eur. J.
765 Wildl. Res. **59**, 271-280.

766

767 Reed, S.E. and Merenlender, A.M., 2011. Effects of management of domestic dogs and
768 recreation on carnivores in protected areas in northern California. Conserv. Biol. **25**, 504-513.

769

770 Ritchie, E.G. and Johnson, C.N., 2009. Predator interactions, mesopredator release and
771 biodiversity conservation. Ecol. Lett. **12**, 982-998.

772

773 Roemer, G.W., Gompper, M.E. and Van Valkenburgh, B., 2009. The ecological role of the
774 mammalian mesocarnivore. Bioscience **59**, 165-173.

775

776 Salo, P., Banks, P.B., Dickman, C.R. and Korpimäki, E., 2010. Predator manipulation
777 experiments: impacts on populations of terrestrial vertebrate prey. Ecol. Monogr. **80**, 531-546.

778

779 Sarmiento, P.B., Cruz, J.P., Eira, C.I. and Fonseca, C., 2010. Habitat selection and abundance
780 of common genets *Genetta genetta* using camera capture-mark-recapture data. Eur. J. Wildl.
781 Res. **56**, 59-66.

782

783 Schowalter, T.D., 1985. Adaptations of Insects to Disturbance, In The Ecology of Natural
784 Disturbance and Patch Dynamics. eds Pickett, S.T.A., and White, P.S. pp. 235-252. Academic
785 Press, New York.

786

787 Schuette, P., Wagner, A.P., Wagner, M.E. and Creel, S., 2013. Occupancy patterns and niche
788 partitioning within a diverse carnivore community exposed to anthropogenic pressures. Biol.

789 Conserv. **158**, 301-312.

790

791 Seymour, C.L. and Dean, W.R.J., 1999. Effects of heavy grazing on invertebrate assemblages
 792 in the Succulent Karoo, South Africa. J. Arid Environ. **43**, 267-286.

793

794 Singleton, G.R., 2010. Rodent Outbreaks: Ecology and Impacts. International Rice Research
 795 Institute, Los Baños.

796

797 Singleton, G.R., Leirs, H., Hinds, L.A. and Zhang, Z., 1999. Ecologically-based management
 798 of rodent pests - re-evaluating our approach to an old problem, In Ecologically-based
 799 Management of Rodent Pests. ed. Singleton, G.R., pp. 17-29. Australian Centre for
 800 International Agricultural Research, Canberra.

801

802 Skinner, J.D. and Chimimba, C.T., 2005. The Mammals of the Southern African Sub-region.
 803 Cambridge University Press, Cambridge.

804

805 Smithers, R.H.N., 1971. The mammals of Botswana. Trustees of the National Museums of
 806 Rhodesia, Salisbury.

807

808 Smithers, R.H.N. and Wilson, V.J., 1979. Check list and atlas of the mammals of Zimbabwe
 809 Rhodesia. Trustees of the National Museums and Monuments, Salisbury.

810

811 Soulé, M.E., Estes, J.A., Miller, B. and Honnold, D.L., 2005. Strongly interacting species:
 812 Conservation policy, management, and ethics. Bioscience **55**, 168-176.

813

814 Stuart, C. and Stuart, T., 2007. Field Guide to the Larger Mammals of Africa. Struik, Cape
 815 Town.

816

817 Suraci, J.P., Clinchy, M. and Zanette, L.Y., 2017. Do large carnivores and mesocarnivores have
 818 redundant impacts on intertidal prey? PLoS One **12**, e0170255.

819

820 Swanepoel, L.H., Swanepoel, C.M., Brown, P.R., Eiseb, S.J., Goodman, S.M., Keith, M.,

Kirsten, F., Leirs, H., Mahlaba, T.a.A.M., Makundi, R.H., Malebane, P., von Maltitz, E.F., Massawe, A.W., Monadjem, A., Mulungu, L.S., Singleton, G.R., Taylor, P.J., Soarimalala, V. and Belmain, S.R., 2017. A systematic review of rodent pest research in Afro-Malagasy small-holder farming systems: Are we asking the right questions? PLoS One **12**, e0174554.

Taylor, P.J., Downs, S., Monadjem, A., Eiseb, S.J., Mulungu, L.S., Massawe, A.W., Mahlaba, T.a.A., Kirsten, F., Von Maltitz, E., Malebane, P., Makundi, R.H., Lamb, J. and Belmain, S.R., 2012. Experimental treatment-control studies of ecologically based rodent management in Africa: balancing conservation and pest management. Wildl. Res. **39**, 51-61.

Tobler, M.W., 2015. Camera base version 1.7 [computer program]. Available from <http://www.atrium-biodiversity.org/tools/camerabase/>.

Vanak, A.T. and Gompper, M.E., 2009. Dogs *Canis familiaris* as carnivores: their role and function in intraguild competition. Mamm. Rev. **39**, 265-283.

Vanak, A.T. and Gompper, M.E., 2010. Interference competition at the landscape level: the effect of free-ranging dogs on a native mesocarnivore. J. Appl. Ecol. **47**, 1225-1232.

Vincent, C., Goettel, M.S. and Lazarovits, G., 2007. Biological control: a global perspective: case studies from around the world. CABI, Cambridge.

Wickham, H., 2016. ggplot2: Elegant Graphics for Data Analysis. Springer, New York.

Williams, S.T., Maree, N., Taylor, P., Belmain, S.R., Keith, M. and Swanepoel, L.H., 2017. Small carnivore ecosystem services data. Figshare [online database]. Available from https://figshare.com/articles/Small_carnivore_ecosystem_services_data/4750807. Accessed 04 December 2017. doi 10.6084/m9.figshare.4750807.

Woodroffe, R., Lindsey, P., Romanach, S., Stein, A. and ole Ranah, S.M., 2005. Livestock predation by endangered African wild dogs (*Lycaon pictus*) in northern Kenya. Bio. Conserv. **124**, 225-234.

854 Zuur, A.F., Ieno, E.N. and Elphick, C.S., 2009. A protocol for data exploration to avoid
855 common statistical problems. *Methods Ecol. Evol.* **1**, 3-14.

856

857